# Gerald Amiel Ballena

# **Professional Summary**

Versatile bioinformatics specialist with focused on general expertise in scalable workflows, computational biology, and machine learning. Skilled in developing scalable pipelines and implementing robust methodologies to tackle complex biological problems.

# Experience

#### 2024 Present

**Project Technical Specialist** 

University of the Philippines, College of Public Health

- Developed scalable bioinformatics workflows for high-throughput sequence analysis.
- O Applied metagenomic profiling techniques to environmental datasets.
- Collaborated with cross-disciplinary teams on projects involving data visualization and computational modeling.

# Education

University of the Philippines Diliman.....

Graduated: July 2022

**Thesis**: In silico assessment of the association of pathogenicity and metal-resistance potential of *Fusarium* spp. Pre-print Link

Accomplishments: DOST ASTHRDP-Scholarship

University of the Philippines Baguio.

Graduated: June 2018

Philippine Science High School CAR Campus.

**Skills** 

Programming.....

Languages: Python, R, Bash, Perl

Installation Toolkits: BiocManager, upcxx

Bioinformatics Tools.

Metagenomics: Kraken2, MetaPhlAn, HUMAnN

Assembly & Binning: MEGAHIT, METAWRAP, CheckM

Genomics: CLC Workbench, Roary, MinHash

Quality Control: FastQC, BUSCO, QUAST Trimming: Trimmomatic, Sickle, Cutadapt

Phylogenetics: RaxML, FastTree, IQ-TREE, phyML, BEAST

Gene Ontology: KEGG, GOseq

Annotation: Prokka, ShortBRED, EggNOG-mapper Others: DESeq2, BBMap Suite, anvio-8, SPAdes Data Analysis & Visualization. **Machine Learning**: scikit-learn, caret (R) Visualization: ggplot2, Plotly, Krona, Shiny, Seaborn, Tableau Technical Skills. Version Control: Git, GitHub **High-Performance Computing:** Slurm Workflow Automation: Snakemake, Conda, YAML (config files) Virtualization: Docker, WSL2 Other Tools..... Technical Writing: Overleaf, TeXStudio Collaboration: Thunderbird, Notion, MS Word (mailing lists) Documentation: TeXStudio, Jupyter Notebooks Certifications and Relevant Coursework Certifications..... AI Fundamentals: Nov 2024 Data Literacy: Nov 2024 Fundamentals Courses. Introduction to Data Engineering: Nov 2024 Intermediate-Level Courses. Advanced-Level Courses. Designing Machine Learning Workflows in Python: ...... Nov 2024 Ongoing Courses. Ensemble Methods in Python: Bagging, Boosting, Stacking Visualizing Geospatial Data in R: Visualizing complex spatial datasets for actionable insights. Azure Fundamentals: Advanced | Score: 180 | Percentile: 99th Data Storytelling: Advanced | Score: 200 | Percentile: 99th

## **Projects**

For more details on public projects, visit GitHub Documentation.pdf.

Note Some repositories including Documentation repo access is currently restricted due to NDA obligations.

### Scripts Repositories.....

Side repo: Scripts for general-purpose tasks, including package management and workflow setup.

Experimental Repository: Exploratory scripts focused on innovative computational techniques.

Health repo: Tools and scripts aimed at health-related data analysis and personal wellness tracking.

Finance repo: WIP; developing scripts for personal finance management.

**Project**4 **repo**: Scripts designed for NGS analyses and workflow automation; details kept private due to project confidentiality.

Kitchen repo: We cookin' some unconventional ideas here.

### Documentation Repositories.

**Documentation repo**: Explanation of every script I've ever written; written in a non-technical tone for non-technical audiences.

**Confidential repo**: LaTeX-compiled reports and documentation of my work as PTS I; restricted access due to confidentiality.

Hunting repo: Includes this CV, my resume, associated . tex files, and certifications; tailored for job hunting.

# Key Projects.

**Bioinformatics** Workflow Development: Designed and implemented modular pipelines for omics data analysis, emphasizing quality control, data assembly, and scalable workflows. Focused on creating reproducible methodologies applicable across diverse datasets.

**Spatial Data Analysis:** Utilized QGIS, R and geospatial libraries to visualize spatial patterns in biological datasets. Developed custom visualization tools for integrating geospatial and genomic data in ecosystem-level studies.

**Public Health Data Exploration**: Developed computational approaches for microbial profiling and identifying markers of interest in public health datasets. Leveraged metagenomic techniques to enable actionable insights for research studies.

#### Development of K-mer Analysis and Statistical Workflow:

- Objective: Build an advanced pipeline for k-mer generation, variance analysis, and distribution modeling.
- O Details: Focused on creating statistical workflows to evaluate parametric and non-parametric fits, automating evaluations with custom Python scripts.
- O Tools & Technologies: Jellyfish, Python, Snakemake, Conda.

# High-Accuracy Sequence Alignment Workflow:

- **Objective:** Create a modular workflow for high-identity sequence alignment with validation across multiple tools.
- O Details: Integrated tools like Bowtie2, BWA, and Minimap2 with standardized outputs to enable comparative analysis.
- O Tools & Technologies: Bowtie2, BWA, Minimap2, KMA, Python, Snakemake.

#### Shannon Entropy Analysis of K-mers and Taxonomic Profiles:

- Objective: Quantify sequence complexity and diversity using entropy-based metrics.
- Details: Developed workflows to calculate and integrate Shannon entropy metrics for assessing genomic diversity and data quality.
- O Tools & Technologies: Jellyfish, Kraken2, Python, Snakemake.

# Comprehensive Workflow for Contaminant Removal and SCG Validation:

- Objective: Design a robust pipeline for contaminant filtering and SCG validation.
- Details: Automated filtering using k-mer-based methods and validated results with SCG retention analysis via BUSCO.
- O Tools & Technologies: BUSCO, Jellyfish, Python, Snakemake, Conda.

#### Pipeline Automation and Dependency Management:

- Objective: Streamline bioinformatics workflows with automated dependency checks and alias creation.
- O Details: Developed:
  - bioconda search.py to identify relevant bioinformatics tools.
  - create aliases.py to automate Bash alias generation.
  - check dependencies.py and append new.py for managing and updating YAML files.
- O Tools & Technologies: Python, YAML, Conda, Bioconda.

#### Version Management and Workflow Validation:

- Objective: Ensure reproducibility in bioinformatics workflows.
- O Details: versioncheck.bash validates installed tool versions against expected configurations, improving workflow consistency.
- O Tools & Technologies: Bash, Python, YAML.

## Ongoing Coursework

Ensemble Methods in Python Visualizing Geospatial Data in R

# Conferences and Symposia

Poster Presentation: 47th PhilSocMicro Conference: Globalization OF Philippine Microbiology: Building Quality Infrastructure

2018

Project Representative: Antimicrobial Resistance (AMR) R&D Scoping and Assessment Launching and Dissemination of Results

# Languages

English: Fluent
Professional proficiency.
Tagalog: Native
Conversational
Ilocano: Can understand
Wernicke's area skill issue.

## Computer Languages (For Fun).....

Python: Fluent

R: Advanced

Best data visualization tool out there (currently).

Bash: Intermediate

Go-to shell scripting language.

Perl: Fair

Would rather "speak" in Bash.

C++: Basic

As fluent as someone who hasn't used it in a decade.

#### References

Available upon request & only if already shortlisted.